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The effect of dietary anthocyanins on biochemical, physiological, and subjective exercise recovery: A systematic review and meta-analysis

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Abstract

Anthocyanins (ACN), the sub-class of (poly)phenols responsible for the red-blue-purple pigmentation of fruit and vegetables, have gained considerable interest in sport and exercise research due to their potential to facilitate exercise recovery. A systematic literature search was performed using PubMed, The Cochrane Library, MEDLINE, SPORTDiscus and CINAHL. Thirty nine studies were included and the standardised mean difference (Hedges *g*) for creatine kinase (CK), anti-oxidative and inflammatory markers, strength, power and delayed onset muscle soreness (DOMS) indices were pooled in separate meta-analyses; meta-regression was also performed on reported ACN dose. Immediately post-exercise there was an increase in antioxidant capacity (*g*: 0.56) and reduced C reactive protein (*g*: -0.24) and tumour necrosis factor α (*g*: -40); P \leq 0.02. Strength was improved with ACN at all time points (*g*: 0.62) and 48 hours (*g*: 0.57) post exercise. The CK was lower 48 hours post-exercise (*g*: -0.31) and there was a trend for a positive association with ACN dose (P = 0.057). This systematic review provides new data showing ACN-rich foods promote functional and subjective recovery likely due to the antioxidant and anti-inflammatory properties of ACN.

1

Introduction

2 Physical exercise places a degree of mechanical and metabolic stress on the body, which both 3 contribute to a common pathological response involving oxidative stress and inflammation (Pyne, 1994). Exercise induced muscle damage (EIMD) is typically characterized by an initial 4 5 insult followed by a secondary inflammatory response, which is more prominent following 6 eccentric actions (Bongiovanni et al., 2020). The EIMD has more immediate implications, 7 including delayed onset muscle soreness (DOMS) and impaired muscular strength and power, which have the ability to compromise performance and quality of training (Clarkson et al., 8 9 1992; MacIntyre et al., 1995). Although the mechanisms and time course can differ, both mechanical and metabolic stress can cause an increase in the appearance of intracellular 10 11 proteins in the blood (e.g. creatine kinase: CK) potentially due to disruptions in calcium homeostasis from a loss of cell membrane integrity (Brancaccio et al., 2007; Tee et al., 2007). 12 In addition, exercise also induces signalling cascades, largely orchestrated by reactive oxygen 13 14 and nitrogen species (RONS), transcriptional release of pro-inflammatory cytokines, (e.g. tumour necrosis factor alpha [TNF- α] and interleukin-6 [IL-6]) and acute phase proteins, e.g. 15 C-reactive protein; CRP (Ebbeling & Clarkson, 1989; Pyne, 1994) and an increase in these 16 immunological markers are thought to be associated with muscle soreness, loss of muscle 17 function as well as overtraining and fatigue (Gleeson et al., 1995; Hecksteden et al., 2016; 18 19 MacIntyre et al., 1995). Given the potential for physiological stress associated with strenuous exercise and the potential for compromised training and/or competition performance due to 20 loss of strength and power and muscle soreness that can last for several days, there has been a 21 22 strong emphasis to identify natural recovery strategies (Bongiovanni et al., 2020; Howatson & van Someren, 2008). 23

Amongst the available strategies, dietary interventions, particularly fruit, have gained considerable attention when it comes to improving recovery following EIMD (Doma et al.,

2020; Naderi et al., 2018). Fruit could influence the recovery process because they contain 26 (poly)phenols which could interact with the secondary cascade associated with EIMD, via their 27 28 antioxidant and anti-inflammatory properties (Bowtell & Kelly, 2019; Pereira Panza et al., 2015). However, not all fruit are equal in terms of (poly)phenolic content and abundance and 29 certain fruit might be more beneficial for exercise recovery. For example, in a simple meta-30 analysis of 25 studies of fruit on recovery from EIMD, berries were reported to have the 31 32 greatest overall effect (Doma et al., 2020). Berries are rich in anthocyanins (ACN), a subclass of (poly)phenols, responsible for the red-blue-purple pigmentation in fruits (Manach et al., 33 34 2004; Pérez-Jiménez et al., 2010). These compounds have gained much research interest due to their propensity to maintain the balance between the oxidative and anti-oxidative systems 35 and reduce inflammatory cytokines (Wang et al., 1999). In vitro ACN have been shown to act 36 37 as more potent antioxidants as compared to some other (poly)phenols (Pojer et al., 2013) and to exhibit anti-inflammatory actions similar or superior to nonsteroidal and other anti-38 inflammatory drugs (Pereira Panza et al., 2015; Seeram et al., 2001). Additionally, ACN might 39 supress pain at both an enzymatic (e.g. cyclooxygenase) and transcriptional (e.g. nuclear factor 40 kappa beta) level (Pojer et al., 2013; Seeram et al., 2001). Moreover, cyanidin-3-glucoside, a 41 common ACN found in berries (Sandoval-Ramírez et al., 2019), has been shown to up-regulate 42 the expression of transcriptional pathways related to muscle function and reduce fatigue in 43 rodent models (Hu et al., 2020; Matsukawa et al., 2017). Other benefits include the potential 44 45 for ACN to enhance blood flow that might aid the removal of waste products and muscle metabolites (Keane et al., 2016b; Rodriguez-Mateos et al., 2019). Recently, Bloedon et al. 46 (2019) reported that ACN-rich whole foods reduce exercise-induced oxidative stress and 47 inflammation, but the effect on muscle soreness and functional (e.g. strength and power) 48 recovery, which are arguably more important measures of exercise recovery (Byrne et al., 49 2004; Damas et al., 2016; Torres et al., 2012), were not reported. Therefore whilst there is 50

some evidence that ACN might be beneficial in facilitating recovery, the narrative nature of these reviews or lack of attention to the potential active compounds or other aspects of recovery means the conclusions are not quantitively derived or based on a complete picture of the available work (Bloedon et al., 2019; Doma et al., 2020; Harty et al., 2019; Naderi et al., 2018; Owens et al., 2019; Vitale et al., 2017). Therefore, the aim of this study was to synthesize and evaluate the effects of anthocyanin-rich foods on biochemical, physiological, and subjective indices of exercise recovery in human trials.

58

Methods

59 *Search strategy*

A systematic literature search of the following electronic bibliographic databases: PubMed and The Cochrane Library as well as searching MEDLINE, SPORTDiscus and CINAHL via EBSCOhost was carried out from inception until August 2020. The search strategy (Supplementary material) was conducted using Medical Subject Heading (MeSH) and Boolean operations devised using two key concepts 1) anthocyanin-rich foods and 2) exercise recovery. Furthermore, the reference list of retrieved literature reviews was hand-searched to find potential articles that could be included in the systematic review.

67 Study selection

The inclusion criteria were as follows: 1) randomised controlled trials; 2) in healthy adult participants (average age ≥ 18 years) regardless of training status; 3) included anthocyanin-rich foods [blackberry, blackcurrant, blueberry, black elderberry, black grape, cherry, chokeberry, rhubarb, strawberry, red wine, plum (Manach et al., 2004; Pérez-Jiménez et al., 2010) or other red-blue-purple berries only where the ACN content was reported] given before exercise (could continue administration after); 4) had a placebo or suitable control; 5) reported haematological markers, functional (e.g. strength or power) or subjective (e.g. visual analogue scales or pain pressure threshold) recovery measures following exercise. For comparability, only similar biomarkers were included in the meta-analysis, these were; creatine kinase (CK) antioxidant (total antioxidant capacity/ status), inflammatory (IL-6, TNF- α or CRP) or oxidative stress (thiobarbituric acid reactive substances; TBARS), antioxidant enzyme activity [superoxide dismutase (SOD) and glutathione peroxidase (GPx)], strength (maximal voluntary contractions; MVC), power (counter movement jumps; CMJ) and visual analogue scales or Likert scales for DOMS.

Exclusion criteria were non-adult, smoker or diseased participants, animal and *in vitro* studies. Studies were also excluded if anthocyanins are given alongside another intervention (i.e. pharmacological agent or dietary supplement; other juice or fruit to increase palatability could be included as long as anthocyanin content was reported) and no appropriate control or reference groups could be identified. Titles and abstracts were independently reviewed by two researchers (RK and CH) to evaluate their eligibility for inclusion in this review. Only full texts that were published in English or had an existing translation were retrieved and examined.

89 Data extraction and quality assessment

90 The study data was extracted into pre-piloted forms by the main reviewer (RK) and checked for accuracy by a second reviewer (KJ). Any discrepancies were resolved by reviewing the 91 original article. The following data was extracted from each study: the first author's last 92 93 name(s), publication date, funding source, participants characteristics, sample size, supplement 94 type, ACN content, dosing strategy and duration, any dietary restrictions, wash out period and 95 type of exercise (metabolic, mechanical or combined], outcome time points, outcome 96 measures, mean \pm SD of the outcomes specified above were also extracted. Where necessary 97 data was extrapolated from figures and graphs and authors were contacted to provide missing data (Abbott et al., 2020; Hurst et al., 2019; Hurst et al., 2020; McCormick et al., 2016; 98

Morehen et al., 2020), if they did not respond within 1 week a follow up was sent, those who 99 did not reply within a month were excluded [e.g. (Beals et al., 2017; Lamb et al., 2019a)] for 100 101 variables where data could not be obtained. A modified PEDro scale (de Morton, 2009) was used to assess the methodological quality of the selected studies. One point could be awarded 102 for each the original 11 items as well as additional items thought to be relevant to the study 103 104 design. Additional criteria were as follows 1) the study acknowledged whether or not they 105 received funding; 2) compliance to the intervention was reported; 3) ACN content was reported 106 in the supplement either according to the manufacturer's nutritional label or confirmed by 107 analysis in the study; participants refrained from taking antioxidant and anti-inflammatory drugs and supplements 4) before and/or 5) during study 6) sample size calculation was 108 included. In studies where a cross-over design an additional item was included '7) a minimum 109 7-day washout between trial treatments'. Thus, a total of 17 points could be awarded for parallel 110 studies and 18 for crossover studies. For parallel studies a score of <7, 7-10, 11-14 and 15-17 111 and for crossover studies <8, 8-11, 12-15 and 16-18 was poor, fair, good and excellent, 112 respectively (Doma et al., 2020). Risk of bias was also assessed according to Cochrane 113 Collaboration guidelines and is represented graphically to indicate the overall quality of all 114 studies (Higgins & Green, 2011). 115

116 Statistical analysis

Standardized mean differences (SMD) were calculated using Hedge's g using independent groups and for parallel studies and paired groups for crossover studies (Borenstein et al., 2019). To calculate the standard deviation within groups for crossover studies the correlation between pairs of observations (r; which was calculated from studies where individual data was provided (Hurst et al., 2019; McCormick et al., 2016)) assumed to be 0.5 (Amiri et al., 2019; Doma et al., 2020; Higgins & Green, 2011) was included. Both study designs were included in an inverse random effects meta-analysis (due to study design heterogeneity) using Stata v.16.0

(StataCorp, College Station, Texas, USA). Initially, studies were sub-grouped by study design 124 to determine whether the inclusion of crossover designs influenced the SMD (Supplemental 125 126 information). Where there were sufficient studies (Jackson & Turner, 2017) separate metaanalyses were conducted for immediately post (\leq 2h), 24 hours post and 48 hours post exercise. 127 For studies that reported measures over several time points, the data were only analysed for the 128 129 most recent in that time interval. Hurst et al (2019) reported different doses so these were 130 pooled to get an overall ES before inclusion in the meta-analysis (Higgins & Green, 2011). If DOMS was measured at different sites, the largest ES was included in the meta-analysis. 131

Secondly, sensitivity analysis was performed by omitting one study at a time to evaluate the 132 potential bias and robustness of the overall SMD. Heterogeneity between studies was 133 determined by the I² statistic. For the I² statistic, I² values $\leq 25\%$, $\leq 50\%$, $\leq 75\%$ and >75%134 indicated no, little, moderate and significant heterogeneity, respectively. To identify potential 135 sources of heterogeneity, moderator analysis was performed using sub-group analysis for 136 137 categorical variables including training status, exercise type and study duration. In addition, where ACN content was reported a meta-regression was conducted on the most reported 138 variables (MVC, DOMS and CK). Potential publication bias for each outcome was evaluated 139 by Egger's test (P < 0.10) and visual inspection of funnel plots (Begg & Mazumdar, 1994). 140 Where publication bias was detected, trim and fill analysis was conducted (Steichen, 2010). 141 The SMD were interpreted as small (>0.2), moderate (>0.5) and (≥ 0.8) large (Sullivan & Feinn, 142 2012) and a Z effect P < 0.05 was considered significant. 143

144

Results

145 Literature search and study characteristics

The search results are presented in Figure 1, following full search and exclusion of irrelevantarticles 39 articles were included in this review. A total of 27 independent group studies and

12 crossover studies with 767 participants were included in this review (Table 1). Of the 148 interventions used tart cherry was the most common (18 studies). Other studies used 149 150 blackcurrant (6 studies), grape (6 studies), blueberry (3 studies), chokeberry (3 studies), bilberry (1 study), plum (1 study) and one a mixed anthocyanin cocktail. The duration of the 151 studies varied greatly with some investigating the acute influence (1-2 h before), most 152 investigating the short-term influence (2-10 days) and some the longer-term influence (20 days 153 154 -8 weeks) of ACN. Most studies were in trained individuals and the median age was 24 (range 18-48) years. The median ACN content, where reported, was 80 (range 8-3600) mg/day. The 155 156 quality of studies was rated as poor (n=1), fair (n=8), good (n=19), excellent (n=11; Table 1). The risk of bias is also represented graphically in Figure 2, which showed the percentage of 157 studies with low, medium and high risk of bias for each domain. The main potential sources of 158 bias came from allocation, blinding of the intervention or did not acknowledged whether they 159 received funding (other bias). 160

161 *The influence of ACN on recovery*

Immediately post exercise there was an increase in TAC (SMD: 0.56; 95% CI: 0.09, 1.03; P = 162 0.02; $I^2 = 61.7\%$) with ACN. ACN also resulted in a moderate reduction in SOD (SMD: -0.42; 163 95% CI: -0.77, -0.07, P = 0.02), TNF- α (SMD: -0.40; 95% CI: -0.72, -0.07, P = 0.02) and a 164 small reduction in CRP (SMD: -0.24; 95% CI: -0.43, -0.06, P = 0.01) at immediately post-165 exercise, with no heterogeneity ($I^2 = 0.0\%$). At 24 hours, SOD remained lower (SMD: -0.46; 166 95% CI: -0.88, -0.03; $I^2 = 16.3\%$) with ACN. Intake of ACN reduced DOMS at 24 hours 167 -0.40, -0.06; P<0.01; I² =0.0%). Strength (MVC) was increased (SMD-0.23; 95% CI: 168 immediately post-exercise (SMD: 0.45; 95% CI: 0.14, 0.75; I² =0.0%), 24 hours post (SMD: 169 0.50; 95% CI: 0.18, 0.82; I²=60.3%) and greatest at 48 hours post (SMD: 0.67; 95% CI: 0.32, 170 1.02; I² =65.4%). At 24 hours power (CMJ) was also increased with ACN (SMD: 0.62; 95% 171 CI: 0.01, 1.24; P=0.047; I²=66.6%). At 48 hours CMJ (SMD: 0.57; 95% CI: 0.04, 1.11; P=0.04; 172

173 $I^2 = 63.9\%$) and CK were lower (SMD: -0.31 95% CI: -0.55, -0.08; P<0.01; $I^2 = 12.8\%$). 174 Sensitivity analysis suggested stable results for these variables (Supplemental material). There 175 was no influence of ACN on any other variables (Figure 3).

176 Subgroup analysis, publication bias and meta-regression

Subgroup analysis is presented in Table 2. Immediately post-exercise ACN increased TAC and 177 decreased SOD, only in metabolically biased exercise. IL-6 was also reduced by metabolic-178 type exercise immediately post and 48 hours post exercise. MVC was improved immediately 179 180 post-exercise with mechanically damaging exercise, whereas 24 hours post and 48 hours post there was a large effect for exercise that had a combined mechanical and metabolic component. 181 182 CRP was lower immediately post exercise, DOMS at 24 hours post exercise and CK at 48 hours 183 post-exercise with combined-type exercise. MVC was improved immediately post-exercise and 24 hours post-exercise in trained and untrained individuals, but only 48 hours post-exercise 184 in trained participants. CRP was lower immediately post-exercise in trained individuals, 185 whereas in the untrained participants IL-6 was lower. 24 hours post-exercise TNFα and IL-6 186 was lower in untrained, whereas trained participants had lower SOD and DOMS at 24 hours 187 and CK at 48 hours. 188

189 Studies of a shorter duration decreased CRP immediately post-exercise and improved MVC and DOMS 24 hours post and CK and CMJ 48 hours post. Studies of a longer duration 190 Increased TAC and reduced SOD, TNFα and CK immediately post. TAC remained higher at 191 48 hours and TBARS and TNFα were reduced in the longer duration studies. There was 192 evidence of publication bias for TBARS (P=0.001) and DOMS (P=0.009) immediately post-193 194 exercise. So too were TAC, CMJ and MVC (P<0.058) 24 and 48 hours post exercise and IL-6 (P=0.004) 48 hours post-exercise. No other publication bias was detected. Trim and fill analysis 195 was done for the above variables resulting in lower DOMS immediately post-exercise (SMD: 196

-0.33; 95% CI: -0.60, -0.06) higher CMJ at 24 hours (SMD: 0.47; 95% CI: 0.12, 0.81) and increased TAC at 48 hours post-exercise (SMD: 0.35; 95% CI: 0.04, 0.67). With trim and fill analysis CMJ was no longer significantly higher at 48 hours (SMD: 0.18; 95% CI: -0.10, 0.47), but there was no other materially different SMDs. Meta-regression suggested a trend for a positive association with ACN dose and CK at 48 hours (P = 0.057; I²=0%), however there was no relationship with MVC or DOMS or CK at any other time point.

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Discussion

The present study represents the most comprehensive picture that synthesizes and evaluates the effects of dietary ACN on exercise recovery from all the available literature including additional analyses that consider ACN dose, exercise type, training status and study duration. These new data showed a beneficial effect for ACN on biochemical, physiological, and subjective recovery following exercise up to and including 48 hours post-exercise.

Dietary intake of ACN resulted in an increase in total antioxidant capacity/status immediately 210 post exercise, which was mirrored by a reduction in SOD at the same point which was still 211 212 reduced 24 hours post exercise, suggesting less reliance on these defence systems over time due to the ability of the ACN to scavenge free radicals (Skarpańska-Steinborn et al., 2006). 213 214 Dietary ACN have antioxidant potential due to the ability for hydrogen (electron) donation and the positively charged oxygen in the flavonoid molecule (Bi et al., 2014). Moreover, the time 215 course aligns with plasma maximum concentrations of ACN and their metabolites, which 216 typically occurs 1-2 hours after ingestion (Hurst et al., 2019; Keane et al., 2016a). In 217 accordance, a number of studies included in this analysis that measured these indices gave an 218 acute dose pre-exercise that would coincide with the peak plasma concentrations (de Lima 219 Tavares Toscano et al., 2019; Hurst et al., 2019; Hurst et al., 2020; Lyall et al., 2009; McAnulty 220

10

et al., 2014; Silvestre et al., 2014). Interestingly, the antioxidant effects of ACN was predominantly seen in exercise with a major metabolic component, which might be attributable to greater exercise-induced oxidative stress owing to higher oxygen consumption during the exercise, whereas a delayed and prolonged generation of RONS after mechanically strenuous eccentric exercise is likely (Fisher-Wellman & Bloomer, 2009), because of the secondary inflammatory-mediated damage that occurs after exercise (Howatson and van Someren, 2008; Owens et al., 2019; Bongiovanni et al., 2020).

The consumption of ACN resulted in reduced CK at 48 hours post, and inflammation (TNFa 228 and CRP) to be reduced immediately post-exercise. As there is an inherent interplay between 229 these markers and RONS (Baird et al., 2012; Lee & Clarkson, 2003), the early antioxidant 230 231 actions of the ACN represents one potential mechanism that might supress the efflux of CK (through reduced cell membrane disruption) and inflammatory indices. The anti-inflammatory 232 properties of ACN are well documented (Fallah et al., 2020; Speer et al., 2020), and might 233 234 relate to their ability to interact with cellular enzymes and signalling pathways (Li et al., 2017). For example, ACN have been shown to reduce inflammatory enzymes such as cyclooxygenase 235 and lipoxygenase (Kirakosyan et al., 2018; Wang et al., 1999), which might be mediated by 236 their ability to inhibit mitogen-activated protein kinase and nuclear factor kappa beta pathways 237 (Pojer et al., 2013). Dependent on exercise modality, intensity and duration CK has been shown 238 to peak 24-72 hours after exercise (Baird et al., 2012). Whereas an acute inflammatory response 239 due to immunological activation typically occurs more rapidly (1-4 hours) and a second wave 240 of inflammation is detectable in a similar timeframe to peak CK (Peake et al., 2017). A lower 241 242 peak in the CK and inflammatory indices might reflect a reduction in muscle damage and also indicate a faster recovery after exercise with ACN compared to a control. While this might 243 relate to the antioxidant capacity of the ACN it could also be because of improved blood flow 244 245 and clearance (Baird et al., 2012; Rodriguez-Mateos et al., 2019). Other meta-analyses have

not suggested an effect of fruit or (poly)phenols on CK (Doma et al., 2020; Hill et al., 2021), 246 it should be acknowledged there are several criticisms of CK as a marker for muscle damage 247 248 especially owing to its high inter and intra-individual variability and its meaningfulness as a recovery index (Brancaccio et al., 2007; Hill et al., 2021; Warren et al., 1999). However, some 249 included studies in the current review found a benefit of ACN-rich foods (Carvalho et al., 2018; 250 251 (Lyall et al., 2009) and an ACN rich cocktail on exercise-induced CK (Lima et al., 2019) it 252 therefore might be that some (poly)phenols such as ACN are more beneficial than others. 253 Moreover, the large number of pooled studies at 48 hours post exercise might account for some 254 of the variability, where the participant numbers amounted to 244.

The aforementioned supports the notion that ACN improves biomarkers related to exercise 255 256 recovery. However, the influence on symptoms such as functional (i.e. strength and power) and muscle soreness indices perhaps are better representations of recovery facilitation and EIMD 257 (Byrne et al., 2004; Damas et al., 2016; Torres et al., 2012). There was an effect of ACN on 258 259 reducing DOMS at 24 hours and recovery of strength loss 0, 24 and 48 hours post exercise, whereas power was only increased 24 and 48 hours post-exercise. Reduced strength loss, and 260 recovery of strength, was greater with ACN, initially for eccentrically biased exercise, but CMJ 261 and MVC were improved 24 hours and 48 hours post-exercise after combined metabolic and 262 mechanically strenuous exercise. Both mechanical and metabolic exercise increase RONS due 263 to mitochondrial oxygen consumption, the increased circulating catecholamines, elevated 264 participation of eccentric muscle contraction-induced damage, inflammatory response and/or 265 the intermittent and repeated sprint actions that can cause temporary ischemic-reperfusion in 266 267 the skeletal muscle (Ascensão et al., 2008; Leeuwenburgh & Heinecke, 2001). Strength loss after exercise has been proposed to be related to oxidative stress (Cakir-Atabek et al., 2019), 268 whereas loss in muscle power might be more synonymous with DOMS and the inflammatory 269 270 response (Byrne et al., 2004). Speculatively, the early increase in antioxidant capacity with

ACN might help to reduce strength loss, whereas the recovery in power coincides with the 271 reduced DOMS at 24 hours post-exercise. These data are of great interest because therapeutic 272 273 recovery interventions (e.g. massage, cold water immersion and compression garments) have shown some benefits in recovery of DOMS, strength and power, but there are limited data to 274 suggest that all facets of recovery can be affected in a positive manner (Brown et al., 2017; 275 276 Davis et al., 2020; Leeder et al., 2012). Whereas, in this review, ACN-rich foods are shown to 277 improve physiological and subjective recovery following strenuous exercise and hence should 278 be an integral consideration for practitioners and exercisers to consider in their diet.

279 Notwithstanding, there are several limitations within the included studies that warrant discussion. Firstly, studies with a crossover study design were included in the meta-analysis 280 281 and these could be influenced by the repeated bout effect (RBE) between experimental trials. The RBE refers to the protective effect afforded by a single bout of eccentric-biased muscle 282 actions that provide a protective effect on subsequent bout of exercise (even if this is performed 283 284 on the contralateral limb) and hence could mask any treatment effect (Howatson & van Someren, 2007). However, including crossover studies did not appear to add to heterogeneity 285 to the results (Figure 3). Secondly, some studies which investigated the effects on functional 286 and subjective recovery after 'real' game play (Abbott et al., 2020; Kupusarevic et al., 2019; 287 Morehen et al., 2020); while these arguably have good application they are heavily confounded 288 by the RBE as well as other recovery practices that might be conducted concurrently. 289 Conversely, some studies used dietary restrictions (Table 1) to reduce phenolic intake. This 290 might lead to an overestimate in the effect, as removal of natural antioxidants from the diet 291 292 might conceptually impair the natural recovery process; therefore ACN might only restore antioxidant capacity whereas the placebo remains in a depleted state. The balance between 293 reducing background noise and ecological validity needs careful consideration in research 294 295 designs (Bowtell & Kelly, 2019). Thirdly, there was a large difference between ACN content

of the interventions and it is not possible to distinguish between different types of ACN, which 296 could have different bioactivities (Rechner & Kroner, 2005). Notwithstanding, ACN content 297 is often reported as cyanidin equivalents (Bell et al., 2016b; Brown et al., 2019; Hutchison et 298 al., 2016; O'Connor et al., 2013) and this compound is an established biomarker of berries 299 (Sandoval-Ramírez et al., 2019) and tart cherries (Seymour et al., 2014) suggesting at least 300 some commonality between the interventions. Moreover, this is the first review to 301 302 comprehensively study ACN on exercise recovery, including a meta-regression of ACN dose. Nonetheless, future studies should try to distinguish the optimum type and dosage of 303 304 anthocyanins for recovery, an important factor highlighted in a recent review (Sabou et al., 2021). Lasty, blinding of studies was a major source of bias, although it is acknowledged that 305 this is an inherent challenge with studies involving functional foods (Brown et al., 2018). 306 307 Therefore, results from this meta-analysis have to be interpreted in light of limitations of the literature highlighted above. Nonetheless, the meta-analytical technique is currently the best 308 method to systematically consolidate evidence from previous work (Haidich, 2010), but it 309 should be conducted with a forensic eye of the literature in order to interpret the information 310 with insight. 311

To summarise, ACN were shown to have an overall beneficial effect on reducing CK, muscle 312 soreness, strength loss and improving power after exercise. This was accompanied by 313 314 attenuated inflammation and increased antioxidant capacity/status following the intake of ACN, suggesting a potential causal link. The information provided by sub-group analyses 315 suggested the most beneficial effect on the biomarkers are following metabolically biased 316 317 exercise and longer-term interventions; whereas shorter duration interventions saw most benefit on physiological variables, which can collectively help inform research designs and 318 application of ACN in exercise recovery. These data provide new information to support the 319

- 320 use of ACN-rich foods in promoting recovery following strenuous exercise that can inform
- 321 exercisers and practitioners.
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- 324

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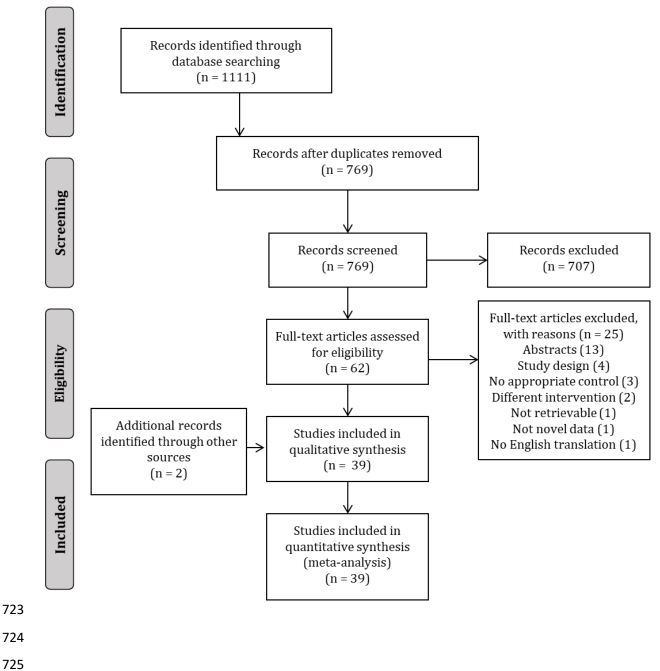


Figure 1. PRISMA flow diagram of included studies: there were 1111 studies identified by the search strategy, 769 non-duplicated records. After screening the titles and abstracts, 62 of the records were deemed potentially eligible for inclusion and full texts were retrieved for further evaluation. Twenty-five articles were excluded and a further 2 found from hand searching, leaving 39 included studies.

Author	or Study Participants design		<i>n</i> (m/f)	Mean age (y)	Supplement	ACN (mg)	Dosing	Exercise	Dietary restrictions [#]	Study quality
Pilaczynska- Szczesniak et al. (2005)	Pll	Well trained male rowers	ACN:9 CTL:10	21 22	Chokeberry juice	3450	3x 50 mL for 4 wks	Incremental rowing test 40% max increased by 10 % each 3 mins to 90%	N	Good
Connolly, McHugh, and Padilla-Zakour (2006)	CO	Male College students	14	22	Tart cherry juice	80	2 x 12oz 8 days (Exercise day 4) 6 day wash out	2 sets of 20 maximal eccentric contractions of the elbow	Ν	Good
Skarpańska- Stejnborn, Basta, and Pilaczyńska- Szcześniak (2006)	Pll	Well trained male rowers	ACN:10 CTL: 9	20 21	Blackcurrant capsules	250 of BC	3x 326 mg capsules for 6 wks	2000m rowing TT	Ν	Fair
Sadowska- Krępa et al. (2008)	Pll	Healthy males	ACN:9 CTL: 5	22 21	Red grape powder	41	3x 390 mg capsules for 6 wks	300m swim test starting at 70- 75% and last 50m at maximal effort	Ν	Poor
Lyall et al. (2009)	CO	Healthy individuals	5/5	48	Black currant capsules	240	4 capsules (2 pre and 2 post exercise) 21 day wash out	30-min row at 80% $\dot{V}O_{2max}$	Ν	Good
Howatson et al. (2010)	Pll	Recreational runners	ACN: 7/3 CTL: 6/4	37 38	Tart cherry juice	80	2 x 236mL for 8 days (Exercise day 6)	Marathon	Ν	Good
Kuehl et al. (2010)	Pll	Recreational runners	ACN: 19/7 CTL: 15/10	38 32	Tart cherry juice	80	2 x 355 mL for 8 days (Exercise day 8)	3 running segments of relay race $(26.3 \pm 2.5 \text{ km})$	Ν	Good
Skarpanska- Stejnborn et al. (2010)	Pll	Well trained male rowers	ACN: 10 CTL: 12	20 21	Black grape extract	38.5	3x 367 mg capsules for 6 wks	Incremental rowing test 40% max increased by 10 % each 3 mins to 90%	Ν	Fair

Table 1. Study characteristics of included studies

Bowtell et al. (2011)	CO	Well-trained male athletes	10	28	Tart cherry juice	547	2 x 30 ml for 10 days (Exercise day 8) 14 day wash out	10 sets of 10 single-leg knee extensions at 80% of their 1RM	Ν	Fair
McAnulty et al. (2011)	Pll	Trained	ACN: 13 CTL: 12	31 33	Blueberries		250g for 6 wks and 375g 1 hour before	2.5h treadmill run	Y	Good
McLeay et al. (McLeay et al., 2012)	CO	Recreationally trained females	10	22	Blueberry smoothie	193	(3 x 200g blueberries on day of exercise)200g each day 2 days after exercise30 day wash out	3 sets of 100 eccentric knee extensions	Y	Fair
O'Connor et al. (2015a)	Pll	Untrained	ACN: 10/10 CTL: 11/9	20 20	Grape powder	8	46g for 45 days	3 sets of 6 reps eccentric elbow flexion at 120% 1RM	Ν	Excellent
Kastello et al. (2014)	CO	Untrained	4/10	21	Tart cherry tablet	200	2x tablet for 20 days (Exercise day 16) 16 day wash out	5 sets of 10 reps of maximal arm contractions	Y	Fair
Bell et al. (2014)	Pll	Well trained male cyclists	ACN: 8 CTL: 8	30	Tart cherry juice	546	2x 30 mL for 7 days (Exercise day 5-7)	High intensity simulated cycling road race 109 min	Y	Excellent
Silvestre et al. (2014)	CO	Male triathletes	6	44	Black grape juice		66 g of grape concentrate 21 day wash out	10 km cycling, 6 km running in sand and 1.5 km swimming at sea	Ν	Fair
Skarpańska- Stejnborn et al. (2014)	Pll	Well trained male rowers	ACN: 10 CTL: 9	21 21	Chokeberry juice	3600	3x 50 mL for 8 wks weeks)	2000m rowing TT	Ν	Good
Bell et al. (2015)	Pll	Well trained male cyclists	ACN: 8 CTL: 8	30	Tart cherry juice	546	2 x 30mL for 8 days (Exercise day 5)	High intensity simulated cycling road race 109 min	Y	Excellent
Levers et al. (2015b)	Pll	Resistance- trained males	ACN: 11 CTL: 12	21 21	Tart cherry capsules	66	1x 480mg capsule for 10 days (Exercise day 8)	10 sets of 10 reps of a barbell back squat at 70% 1-RM	Ν	Excellent

Toscano et al. (2015)	Pll	Recreational runners	ACN: 11/4 CTL: 11/2	43 36	Grape juice	53 mg/L	2 x 5ml/kg for 28 days (Exercise day 26)	Time to exhaustion at anaerobic threshold	Ν	Good
Bell et al. (2016a)	Pll	Trained soccer players	ACN: 8 CTL: 8	25	Tart cherry juice	73.5	2 x 20 mL for 7 days (Exercise day 5)	Adapted LIST 90 min	Y	Good
Hutchison et al. (2016)	Pll	Untrained	ACN: 1/7 CTL: 2/6	20 21	Blackcurrant nectar	151.4	2 x 455 mL for 8 days (Exercise on day 5)	3 sets of 10 reps of eccentric contractions using a bar weighted with 115% of 1RM	Ν	Good
Levers et al. (2016)	Pll	Endurance trained runners	ACN: 8/3 CTL:10/ 6	21 22	Tart cherry capsules	66	1 capsule for 10 days (Exercise day 8)	Half marathon	Ν	Excellent
McCormick et al. (2016)	CO	Well trained male water-polo players	9	19	Tart cherry juice	819	90 ml for 6 days 35 day wash out	Simulated water polo match 60 min	Ν	Good
Petrovic et al. (2016)	Pll	Male handball players	ACN:8 CTL: 7	19 18	Chokeberry juice		1x 100 mL of chokeberry juice for 4 wks	Training camp: a combination of aerobic, strength and conditioning twice per day, lasting 3 h in total	Ν	Good
Beals et al. (2017)	Pll	Recreationally active	ACN: 9/6 CTL: 10/4	wks 26 Tart cherry 64 2x 30g for 12 days 25 powder (Exercise day 5)		Repetitive, maximal effort isokinetic concentric/eccentric contractions of the quadriceps until the fatigue	Ν	Excellent		
Lynn et al. (2018)	Pll	Recreationally trained runners	ACN: 8/3 CTL: 8/2	31 31	Bilberry juice	80	2 x 200 mL for 8 days (Exercise day 6)	Half marathon	Ν	Good
Assunção Carvalho et al. (2018)	Pll	Well trained male handball players	ACN:12 CTL: 13	19	Plum nectar	53.5	2x 5mL/kg for 28 days	Training camp: 3x 60min sessions of general strength and moderate intensity endurance 2x maximal power and speed sessions and 5 x strength and skill sessions a week for 4 weeks	Ν	Good

Brandenburg and Giles (2019)	CO	Recreational runners	24	31	Blueberry powder		3 x 24g for 4 days (Exercise day 4) 10-14 day wash out	8km treadmill time trial	Y	Good
Brown, Stevenson, and Howatson (2019)	Pll	Trained female dancers	ACN: 10 CTL: 10	19	Tart cherry juice	73.5	2x 30 mL for 8 days (Exercise day 5)	15×30 m maximal sprints with a rapid 10 m deceleration phase	Ν	Good
de Lima Tavares Toscano et al. (2019)	СО	Recreationally trained runners	14	39	Purple grape juice		10 ml/kg (2h before exercise) 7 day wash out	Treadmill running test at 80% of their $\dot{V}O_{2max}$ until exhaustion.	Ν	Excellent
Hurst et al. (2019)	Pll	Healthy individuals	ACN: 8 CTL: 8	44 42	Blackcurrant extract	58.4	0.8mg/kg 1h before exercise	30min row at their predicted 70% $\dot{V}O_{2max}$	Y	Good
Hurst et al. (2019)	Pll	Healthy individuals	ACN: 8 CTL: 8	44 42	Blackcurrant extract	131.2	1.6 mg/kg 1h before exercise	30min row at their predicted 70% $\dot{V}O_{2max}$	Y	Good
Hurst et al. (2019)	Pll	Healthy individuals	ACN: 8 CTL: 8	44 42	Blackcurrant extract	240	3.2 mg/kg 1h before exercise	30min row at their predicted 70% $\dot{V}O_{2max}$	Y	Good
Kupusarevic, McShane, and Clifford (2019)	CO	Well-trained male rugby union players	10	28	Tart cherry juice		2 x 30 mL for 5 days (Exercise day 3) Wash out not reported	Competitive rugby union match	Ν	Fair
Lamb et al. (2019b)	Pll	Untrained males	ACN: 12 CTL: 12	24 24	Tart cherry juice	8	2 x 30 mL for 5 days (Exercise day 5)	5 sets of 10 reps maximal voluntary eccentric contractions of the elbow flexor	Y	Excellent
Lima et al. (2019)	Pll	Untrained males	ACN: 15 CTL: 15	22 23	Mixed plum, blueberry, maquiberry, raspberry and cranberry concentrates	58	2 x 240mL 8 days (Exercise day 5)	running downhill (-15%) for 30 min at 70% of their VO _{2max}	Ν	Good

Quinlan and Hill (2020)	Pll	Team-sport players	ACN: 4/6 CTL: 4/6	28 25	Tart cherry juice		2x 30 ml for 8 days (Exercise day 6)	LIST	Ν	Fair
Abbott et al. (2020)	CO	Well trained male soccer players	10	19	Tart cherry juice		2 x 30 mL for 3 days (Exercise day 1) 14-28 day wash out	Competitive soccer match	Y	Good
Costello et al. (2020)	Pll	Recreational runners	ACN: 6/4 CTL: 6/4	30 29	Blackcurrant extract	110	2x capsules for 10 days (Exercise day 8)	Half marathon	Ν	Excellent
Hurst et al. (2020)	Pll	Healthy individuals	ACN: 17 CTL: 17	38 38	Blackcurrant extract	240	2x capsules for 5 weeks	30min row at their predicted 70% $\dot{V}O_{2max}$	Y	Excellent
Morehen et al. (2020)	СО	Well trained male rugby league players	11	18	Tart cherry juice	640	2x 30mL for 7 days (Exercise day 6) 5 day wash-out	Competitive rugby league match	Ν	Good

Table 1: Abbreviations: anthocyanin (ACN); Blackcurrant (BC); Cross-over (C); Loughborough intermittent shuttle test (LIST); Parallel (Pll), Repetitions (reps) [#] apart from exclusion of the product under investigation.

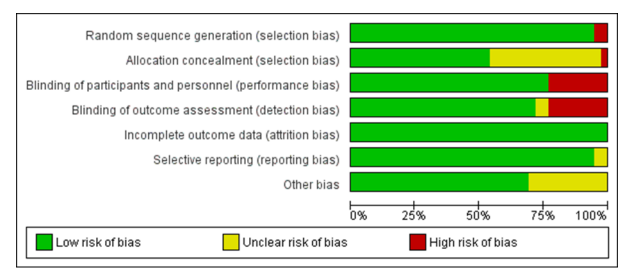


Figure 2. Risk of bias of included studies.

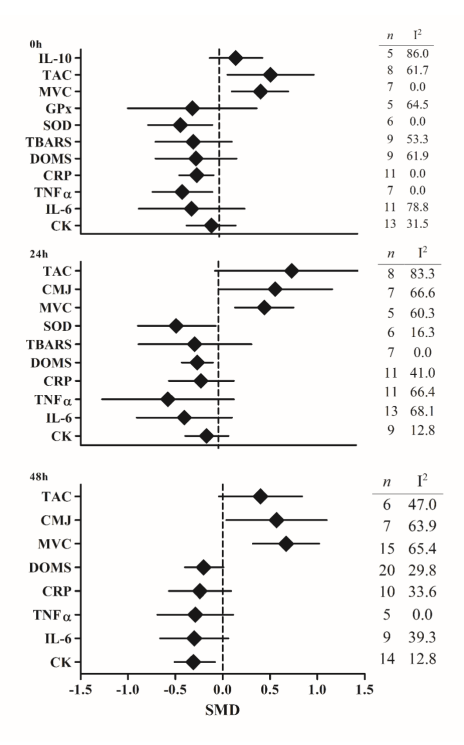


Figure 3. Summary forest plot of findings for anthocyanin (ACN) intake on exercise recovery relative to a control immediately post (top), 24 hours post (middle) and 48 hours post (bottom) exercise. Data missing for timepoint if less than 5 studies (N/A). For maximal voluntary contraction (MVC), countermovement jump (CMJ) and total antioxidant capacity/status (TAC) right side favours ACN. For interleukin 6 (IL-6), tumour necrosis factor alpha (TNF α), C-reactive protein (CRP), thiobarbituric acid reactive substances (TBARS), creatine kinase (CK), superoxide dismutase (SOD) and glutathione peroxidase (GPx), and delayed onset of muscle damage (DOMS) left side favours ACN. (*n* = number of studies, I² statistic for heterogeneity).

				Exercise type	0				Subgr		Stu	Study duration				
Variable		Metabolic	п	Mechanical	$\frac{e}{n}$	Combined	n	Trained	Training status Trained <i>n</i> Untrained <i>n</i>				uy ui n	Long	n	
v al lable		Wetabolic	n	Wiechanical	п	Immedi			п	Olitianieu	n	Short	п	Long		
IL10	SMD (95%CI) I ²	0.72 (0.01, 1.4) N/A	1	0.42 (-0.40, 1.24) N/A	1	0.76 (-0.18, 1.71) 92.7	3	0.82 (-0.4. 2.1) 89.1	4	0.76 (-0.2, 1.7) N/a	1	0.01 (-0.4, 0.4) 0.0	3	2.15 (-0.8, 5.1) 93.1	2	
TAC	SMD (95%CI) I ²	1.07 (0.4, 1.8) 60.6	4	0.10 (-0.4, 0.6) 0.0	2	0.13 (-0.6, 0.9) 36.6	2	0.56 (0.1, 1.0) 61.7	8			0.28 (-0.1, 0.6) 20.0	5	1.23 (0.2, 2.3) 71.3	3	
MVC	SMD (95%CI) I ²			0.47 (0.1, 0.9) 0.0	4	0.41 (-0.1, 0.9) 0.0	3	0.31 (-0.1, 0.7) 0.0	5	0.78 (0.2, 1.3) 0.0	2	0.45 (0.1, 0.8) 0.0	7			
GPX	SMD (95%CI) I ²	-0.40 (-1.3, 0.5) 72.1	4	0.04 (-0.8, 0.9) N/A	1			-0.37 (-1.2, 0.5) 73.0	4	0.00 (-1.1, 1.1) N/A	1	0.04 (-0.8, 0.9) N/A	1	-0.40 (-1.3, 0.5) 72.1	4	
SOD	SMD (95%CI) I ²	- 0.63 (- 1.2, -0.1) 0.0	3	-0.13 (-0.7, 0.44) 0.0	2	-0.54 (-1.4, 0.3) 0.0	1	- 0.42 (- 0.8 , - 0.1) 0.0	6			-0.26 (-0.7, 0.2) 0.0	3	- 0.63 (- 1.2, -0.1) 0.0	3	
DOMS	SMD (95%CI) I ²			-0.04 (-1.2, 1.1) 73.9	2	-0.31 (-0.8, 0.2) 62.2	7	-0.35 (-0.8, 0.1) 56.5	8	0.55 (-0.3, 1.4) N/A	1	-0.25 (-0.7, 0.2)	9			
TBARS	SMD (95%CI) I ²	-0.79 (-1.7, 0.1) 62.6	3	-0.14 (-0.6, 0.3) 0.9	4	0.12 (-0.7, 0.9) 60.4	2	-0.40 (-0.8, 0.00) 35.6	8	0.47 (-0.1, 1.0) N/A	1	-0.51 (-1.3, 0.3) 74.2	5	-0.08 (-0.5, 0.4) 0.0	4	
CRP	SMD (95%CI) I ²	-0.20 (-0.6, 0.2) 0.0	2	-0.25 (-0.7, 0.2) 0.0	4	- 0.26 (- 0.5 , - 0.02) 0.0	5	-0.24 (-0.4, -0.1) 0.0	9	-0.24 (-0.7, 0.2) 0.0	2	-0.24 (-0.4, -0.1) 0.0	9	-0.23 (-0.7, 0.2) 0.0	2	
ГНБа	SMD (95%CI)	-0.38 (-0.8, 0.1)	4	-0.45 (-1.3, 0.4)	1	-0.48 (-1.4, 0.5)	2	-0.34 (-0.7, 0.03)	6	-0.60 (-1.3, 0.1)	1	-0.29 (-0.7, 0.1)	5	-0.59 (-1.1, - 0.04)	2	
	I^2	0.0		N/A		53.5		0.0		N/A		0.0		0.0		
IL-6	SMD (95%CI) I ²	-0.88 (-1.4, -0.4) 6.6	4	-0.05 (-0.9, 0.8) N/A	1	0.03 (-0.9, 1.0) 85.5	6	-0.19 (-0.8, 0.4) 77.6	10	-1.24 (-2.0, -0.5) N/A	1	-0.44 (-2.0, 0.1) 61.9	8	0.20 (-1.7, 2.1) 92.8	3	
CK	SMD (95%CI)	-0.16 (-0.6, 0.3)	6	0.25 (-1.0, 1.5)	2	-0.12 (-0.4, 0.2)	6	-0.01 (-0.3, 0.3)	12	-0.70 (-1.7, 0.3)	2	-0.00 (-0.3, 0.3)	12	-0.83 (-1.6, - 0.04)	2	
	I^2	38.2		78.1		0.7 24h pos		22.2		52.4		19.3		11.7		
		151(04.25)		0.55 (1.2. 2.4)			n pos			1.55 (0.5.2.6)		0.04 (0.6, 0.5)		1 5 (0 1 0 0)		
ГАС	SMD (95%CI) I ²	1.51 (-0.4, 3.5) 90.2	3	0.56 (-1.3, 2.4) 88.6	2	0.16 (-0.6, 0.9) 38.6	2	0.68 (-0.2, 1.6) 83.9	6	1.55 (0.5, 2.6) N/A	1	-0.04 (-0.6, 0.5) 27.5	3	1.5 (0.1, 2.9) 85.9	3	
CMJ	SMD (95%CI) I ²					0.62 (0.01, 1.2) 63.9	6	0.62 (0.01, 1.2) 63.9	6			0.62 (0.01, 1.2) 63.9	6			
MVC	SMD (95%CI) I ²	0.97 (-0.1, 2.0) N/A	1	0.22 (-0.2, 0.6) 61.6	9	0.99 (0.6, 1.4) 0.0	5	0.58 (0.1, 1.1) 66.8	9	0.43 (0.02, 0.9) 55.3	6	0.55 (0.2, 0.9) 65.4	13	0.30 (-0.1, 0.7) 0.0	2	
SOD	SMD (95%CI) I ²	-0.42 (-1.0, 0.1) 34.8	4			-0.60 (-1.4, 0.2) N/A	1	-0.46 (-0.9, -0.03)	5			-0.60 (-1.4, 0.2) N/A	1	-0.42 (-1.0, 0.1) 34.8	4	
DOMS	SMD (95%CI)	-0.25 (-1.0, 0.4)	2	-0.14 (0.4, 0.1)	8	-0.33 (-0.6, -0.08)	10	-0.33 (-0.5, -0.1)	13	-0.09 (-0.4, 0.2)	7	-0.25 (-0.4, - 0.07)	18	-0.14 (-0.5, 0.3)	2	
	\mathbf{I}^2	0.0		15.0		0.0		0.0		2.1		0.0		0.0		
ΓBARS	SMD (95%CI)	-0.46 (-1.3, 0.3)	5	0.27 (-0.3, 0.9)	2			-0.41 (-1.0, 0.2)	6	0.55 (-0.03, 1.1)	1	0.41 (-0.00, 0.9)	3	-0.74 (-1.5, - 0.02)	4	
CRP	I ² SMD (95%CI) I ²	83.5 -0.26 (-0.7, 0.2) 0.0	3	0.0 -0.17 (-0.8, 0.5) N/A	1	-0.19 (-0.9, 0.5) 67.6	5	73.9 -0.16 (-0.6, 0.3) 46.8	8	N/A -0.35 (-0.9, 0.2) N/A	1	0.0 -0.22 (-0.6, 0.2) 49.9	7	69.9 -0.03 (-0.8, 0.7) 25.3	2	

Table 2. Subgroup analysis on moderator variables for effect of ACN on recovery

TNFα IL-6	SMD (95% CI) I ² SMD (95% CI)	-0.34 (-0.9, 0.2) 0.0 0.15 (-0.3, 0.7)	3	-2.2 (-3.4, -1) N/A -1.25 (-3.1, 0.6)	1	0.12 (-0.7, 0.9) N/A -0.41 (-1.2, 0.4)	1	-0.19 (-0.6, 0.3) 0.0 -0.17 (-0.6, 0.3)	4	-2.2 (-3.4, -1) N/A -2.26 (-3.4, -1.1)	1	0.10 (-0.5, 0.7) 0.0 -0.04 (-0.5, 0.4)	2	-1.03 (-2.0, - 0.01) 67.9 -1.18 (-2.5, 0.1)	3
CK	I ² SMD (95%CI) I ²	19.0 -0.17 (-0.5, 0.2) 0.0	5 6	86.0 0.38 (-0.4, 1.1) 48.0	2	63.3 -0.32 (-0.7, 0.1) 0.0	4 5	48.4 -0.09 (-0.3, 0.2) 0.0	8 12	N/A -0.42 (-1.1, 0.3) N/A	1	40.9 -0.11 (-0.4, 0.2) 8.4	9	78.6 -0.18 (-0.6, 0.3) 0.0	4
48h post															
TAC	SMD (95%CI) I ²	0.35 (-0.4, 1.1) N/A	1	-0.04 (-0.5, 0.5) 0.0	2	0.78 (0.03, 1.5) 56.9	3	0.40 (-0.04, 0.9) 79.4	6			0.15 (-0.2, 0.5) 0.0	1	0.94 (-0.3, 2.2) 76.1	2
СМЈ	SMD (95%CI) I ²					0.57 (0.04, 1.1) 63.9	6	0.57 (0.04, 1.1) 63.9	6			0.71 (0.1, 1.3) 63.5	6	-0.20 (-1.0, 0.6) N/A	1
MVC	SMD (95%CI) I ²	1.26 (0.2, 2.4) N/A	1	0.39 (-0.00, 0.8) 63.4	9	1.19 (0.6, 1.8) 53.1	5	0.83 (0.4, 1.2) 50.1	9	0.44 (-0.2, 1.0) 76.7	6	0.78 (0.4, 1.2) 65.5	13	0.12 (-0.3, 0.5) 0.0	2
DOMS	SMD (95%CI) I ²	-0.28 (-1.0, 0.4) 0.0	2	-018 (-0.6, 0.2) 55.8	8	-0.20 (-0.5, 0.1) 19.2	10	-0.18 (-0.4, 0.1) 10.6	13	-0.19 (-0.6, 0.2) 55.9	7	-0.20 (-0.4, 0.03) 34.3	18	-0.20 (-0.7, 0.3) 16.2	2
CRP	SMD (95%CI) I ²	-0.10 (-0.8, 0.6) 46.2	3	-0.30 (-0.8, 0.2) 0.0	2	-0.32 (-0.9, 0.3) 54.7	5	-0.22 (-0.6, 0.2) 39.0	9	-0.42 (-1.0, 0.2) N/A	1	-0.30 (-0.7, 0.1) 37.2	8	-0.09 (-0.8, 0.6) 53.7	2
TNFα	SMD (95%CI) I ²	-0.08 (-0.8, 0.6) 0.0	2	-0.49, (-1.3, 0.3) 0.0	1	-0.38 (-1.1, 0.4) 26.5	2	-0.29 (-0.7, 0.1) 0.0	6			-0.29 (-0.7, 0.1) 0.0			
IL-6	SMD (95%CI) I ²	- 0.70 (- 1.3 , - 0.1) 0.0	3	-0.16 (-0.6, 0.3) 0.0	2	-0.21 (-0.9, 0.5) 64.0	4	-0.28 (-0.7, 0.1) 44.7	8	-0.58 (-1.6, 0.4) N/A	1	-0.30 (-0.7, 0.1) 39.3	9		
СК	SMD (95%CI)	-0.33 (-1.3, 0.6)	3	-0.24 (-0.7, 0.3)	4	-0.36 (-0.7, -0.06)	7	-0.27 (-0.5, -0.01)	12	-0.60 (-1.2, -0.0)	2	-0.29 (-0.5, - 0.04)	13	-0.58 (-1.4, 0.2)	1
-	\mathbf{I}^2	60.1	-	41.8	-	0.0		17.4		0.0		16.8	-	N/A	

Subgroup analysis for: countermovement jump (CMJ); C-reactive protein (CRP); creatine kinase (CK); delayed onset of muscle damage (DOMS); glutathione peroxidase (GPx); interleukin (IL-); maximal voluntary contraction (MVC); superoxide dismutase (SOD); thiobarbituric acid reactive substances (TBARS); total antioxidant capacity/status (TAC) and tumour necrosis factor alpha (TNF α). Grey cells indicate no studies, *n*= number of studies.

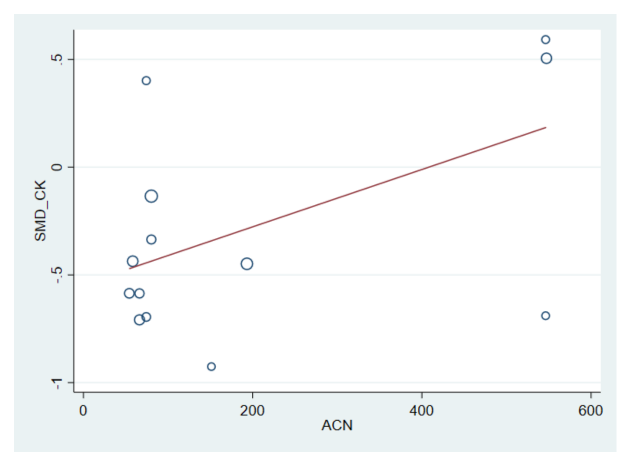


Figure 4. Bubble plot with fitted meta-regression for creatine kinase (CK) at 48h post exercise.